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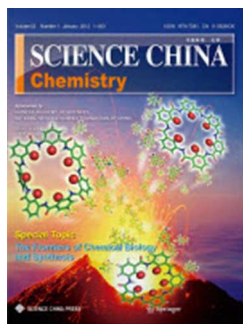
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**Comparison of reduced point charge models of proteins:
Molecular Dynamics simulations of Ubiquitin**

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Abstract

Reduced point charge models of amino acids are used to model Ubiquitin (PDB: 1UBQ). They are designed (i) from local extremum positions in charge density (CD) distribution functions built from the Poisson equation applied to smoothed molecular electrostatic potential functions, or (ii) from local maximum positions in promolecular electron density distribution (ED) functions. Charge values are fitted *versus* all-atom Amber99 molecular electrostatic potentials. The program GROMACS is used to generate molecular dynamics trajectories of the protein, under various implementation schemes, solvation, and temperature conditions. Point charges that are not located on atoms are considered as virtual sites with a null mass and radius. The results illustrate that secondary structure is best preserved with the CD-based model at low temperatures and in vacuum. This indicates that local potential energy wells are consistent with the all-atom model. However, at room temperature, the structure is best conserved when point charges are forced to be located on atoms, due to a better description of the Coulomb 1-4 energy terms. The ED-based model, generated at a lower resolution, led to the largest discrepancies *versus* the all-atom case. The CD-based model allows the formation of protein-water H-bonds with geometrical properties similar to the all-atom ones. Contrarily, intra-molecular H-bonds are not well described. Structural, thermodynamical, and dynamical properties of proteins modelled with reduced point charge models are also significantly affected by the choice of the solvent force field.

Keywords

Molecular electrostatic potential, electron density, smoothing of molecular fields, critical points, point charge model, protein, Ubiquitin

1 Introduction

Topological analyses, also known as critical point (CP) analyses, of three-dimensional (3D) molecular properties such as electron density (ED) distribution functions are a well-known subject of research [1]. For example, the topology of an ED distribution function is particularly useful to interpret bonding and interactions through the Bader's Atom-In-Molecule theory [2-5] with applications to various chemical systems [6-8], in crystallography [9-13] and particularly in H-bond analysis [14,15], and in molecular similarity analysis [16-19].

Intermolecular interactions can also be analysed through the topography of molecular electrostatic potential (MEP) functions. Calculations have initially been proposed by Gadre and coll. [20,21] and later by Leboeuf *et al.* [22] who showed how MEP CPs are related to the electronic structure (π bonds, lone pairs, ...) of the investigated molecules. It was also illustrated that promolecular representations, based on non-interacting spherical atoms, nevertheless provide acceptable results for the analysis of MEP minima along internuclear axes [23,24]. Mata *et al.* [25] reported that zero-flux surfaces occurring in a MEP of an isolated molecule are also observed between atoms, like for the ED, but the actual partition of the space in volumes is different from that of the ED. The authors associated each local maxima (nuclei) and minima with electrophilic and nucleophilic sites, respectively, with the corresponding basins indicating their influence zones [25]. They gave a particular application to H bonds in ref. [26]. By iteratively visualizing the direction of the MEP gradient calculated on spherical surfaces centered at various molecular points, Roy *et al.* [27] determined the critical points of the MEP function, and proposed that the recognition between two species occurs when the geometry is such that a change in the nature of their MEP CPs is observed [28]. Yeole *et al.* also proposed methods to map the CPs of ED and MEP at any level of QM theory [29,30].

Rather than working at atomic resolution, we focus on the analysis of smoothed molecular properties. In recent publications, we described reduced point charge models (RPCMs) [31,32] built from CP analyses of smoothed molecular properties, and their particular applications to Molecular Dynamics (MD) simulations of proteins [33,34].

In the last paper [34], we specifically focussed on the implementation of the models using the program GROMACS [35,36] which allows to describe non-atomic charges as virtual sites with a null mass, connected to the molecular structure through geometrical rules. A detailed description of the reduced point charge templates and representations of the amino acid (AA) was achieved, with a particular application to MD simulations of the Ubiquitin-UIP1 complex. The analysis focussed on the comparison of energetic (Lennard-Jones, short-range Coulomb), structural (secondary structure, gyration radius, H-bond geometry), and dynamical (atom fluctuations, self-diffusion coefficient) results. Two 3D molecular properties, the MEP and the promolecular ED leading to different RPCMs, were considered. For the first case [31,32], a limited number of point charges was obtained through the search for the maxima and minima of a smoothed version of the charge density (CD) generated by the atomic charges defined in Amber99 (or Amber99SB) FF [37]. For the second case, the point charges

were obtained through a search of the maxima of the full promolecular ED of the molecular structure and, as in the first approach, charge values were assigned to those maxima using a least-square charge fitting procedure. This last molecular property is easily calculated using the so-called Promolecular Atom Shell Approximation (PASA) formalism that was developed by Amat and Carbó-Dorca [38,39]. Those two approaches led to various implementation schemes which are also applied in the present paper.

In the present work, we carry on with the assessment of the RPCMs briefly described hereabove with an additional focus on free energy differences between the various point charge models, on H-bond and water dynamics, and on the effect of the water force field (FF). Various solvation and temperature conditions are considered. The selected system is Ubiquitin (PDB: 1UBQ), a protein that is largely studied in literature. At this stage of our research work, only the electrostatic part of the FF is modified. Energetic, structural, and dynamical properties are calculated and compared to the all-atom ones, which is easily achieved as no conversion stage is required between the reduced and all-atom models.

2 Computational methods

The mathematical formalism that was used to design a molecular reduced point charge representation of a protein and its corresponding charge values is briefly presented. Additional details are available in refs. [31,32].

2.1 Critical points of a smoothed molecular property

Based on the formalism presented by Kostrowicki *et al.* [40], a smoothed version of the PASA ED model is calculated as a sum over individual atomic ED distribution functions, $\rho_{a,s}(r)$:

$$\rho_{a,s}(r) = \sum_{i=1}^3 \alpha_{a,i} e^{-\beta_{a,i} r^2} \quad (1)$$

with:

$$\alpha_{a,i} = Z_a w_{a,i} \left(\frac{2\zeta_{a,i}}{\pi} \right)^{3/2} \frac{1}{(1 + 8\zeta_{a,i}s)^{3/2}} \text{ and } \beta_{a,i} = \frac{2\zeta_{a,i}}{(1 + 8\zeta_{a,i}s)} \quad (2)$$

where Z_a , $w_{a,i}$ and $\zeta_{a,i}$ are the atomic number of atom a , and the two fitted parameters, respectively. The smoothing factor s is given in bohr² (1 bohr = 0.52918 10⁻¹⁰ m).

The smoothed analytical CD distribution function $\rho'_{a,s}(r)$ that is obtained from an atomic charge q_a and the Poisson equation can be expressed as:

$$\rho'_{a,s}(r) = \frac{q_a}{(4\pi s)^{3/2}} e^{-r^2/4s} \quad (3)$$

where a and r stand for the atom index and the distance *versus* the atom position, respectively. Unsmoothed functions are obtained by setting $s = 0$ bohr².

To locate CPs, an algorithm initially described by Leung *et al.* [41] was implemented to follow the trajectories of the CPs, more specifically, the maxima and/or minima in a CD or ED function, as a function of the degree of smoothing. At scale $s = 0$, each atom of a molecular structure is considered as the starting point of a trajectory that is generated as s increases. Trajectories are merged when their inter-distance is lower than a predetermined value (SI 1). It has the effect to progressively reduce the number of extrema in the 3D molecular property.

2.2 Design of amino acid reduced point charge models

CD-based templates

The design of the AA point charge templates was achieved in four stages. First, isolated AA structures were assigned Amber99 atom charges using PDB2PQR [42]. Side chain extrema were located using our merging/clustering algorithm applied to the CD distribution functions smoothed at $s = 1.7$ bohr² (SI 1). This was carried out separately for the positively and negatively charged atoms. Second, the charge values of the resulting peaks and pits considered together were fitted *versus* the all-atom MEP generated from the side chain atoms only. In this procedure, several rotamer descriptions were taken into account according to their occurrence probability (see Table 2 of [31]). Third, the main chain point charges were located in accordance with the motif found for the central Gly residue in an extended poly-

Gly strand [31] and, fourth, a second charge fitting procedure (SI 1), now carried out *versus* the MEP calculated using all the AA atoms, was applied to determine the charge values of the main chain point charges while preserving the side chain point charges first obtained. All main chain point charges, observed to be located very close to the C and O atoms, were set exactly on those atoms. The AA models, that consist of two point charges for the main chain and up to six point charges for the side chain, are detailed in ref. [34]. In the further parts of the present paper, the model will be referred to as model mCD (Figure 1).

A second point charge description was derived from the model described above. In this second model, to fully facilitate the implementation of the AA models in GROMACS [35,36], most of the point charges were set exactly on atoms of the residues, and a charge fitting algorithm was again applied (SI 1). Results are presented in ref. [34]. The model will be referred to as model mCDa further in the text (Figure 1).

PASA-based templates

CP searches of the PASA ED distribution functions were carried out to generate even coarser charge descriptions for the AAs. Indeed, with the CD distribution functions depicted above, it is not possible to obtain less than two main chain point charges per residue, *i.e.*, one negative and one positive charge associated with the O and C atoms, respectively. Within the framework of the PASA, the ED depends only on the atomic number Z_a of the atoms, not on their charge (Eqs. 1 and 2). In the further parts of this paper, the model, that consists of one point charge on the main chain and no more than two charges on the side chain, will be referred to as model mPASA (Figure 1). Its implementation within the program GROMACS is detailed in ref. [34].

Automated point charge generation procedure

The point charge templates described above are established for isolated AA structures. Their properties are thus independent on the neighbourhood of a particular protein. To study large proteins, an automation stage was developed to rapidly locate point charges on the whole structure. It is fully based on the application of a superimposition algorithm of CP templates of each AA onto their corresponding all-atom structure of the protein under study. We used the program QUATFIT [43] to, first, superimpose a limited set of atoms from the template on the studied structure, and then use the resulting transformation matrix to generate the corresponding point charge coordinates. The GROMACS topology file, wherein point

charges are defined as virtual sites, is further generated through an in-house program that outputs geometrical parameters as reported and discussed in [ref.](#) [34].

2.3 Free energy calculations

In the present paper, we used two methods to evaluate the free energy difference between the all-atom and RPCMs. The first method belongs to the so-called thermodynamical integration (TI) approaches where the difference in configurational free energy between two states 0 and 1, $\Delta F_{0 \rightarrow 1}$ (or $\Delta G_{0 \rightarrow 1}$ at constant pressure), is calculated using a numerical integration:

$$\Delta F_{0 \rightarrow 1} = \int_0^1 \left\langle \frac{\partial U(\lambda)}{\partial \lambda} \right\rangle_{\lambda} d\lambda \quad (4)$$

where the brackets $\langle \rangle_{\lambda}$ stand for ensemble averages calculated over a MD simulation carried out with the potential energy function $U(\lambda)$ defined as:

$$U(\lambda) = \lambda U_1 + (1 - \lambda) U_0 \quad (5)$$

In the particular slow-growth approach [44], λ is progressively brought from 0 to 1 along the isothermal MD trajectory. Even if the approach is known to be debatable, especially when states 0 and 1 largely differ, *i.e.*, when the transformation cannot be considered as reversible, it has the advantage to require only one MD simulation to obtain an estimate of $\Delta F_{0 \rightarrow 1}$. The second approach, known as the Bennett's acceptance ratio (BAR) [45,46], is based on the statistical mechanics expression (SI 1):

$$\Delta F_{0 \rightarrow 1} = \beta^{-1} \ln \frac{Z_0}{Z_1} \quad (6)$$

where $\beta^{-1} = kT$. Z_i is the configurational integral, *i.e.*, the contribution to the partition function that involves the interaction energy U_i between the particles of a system. The method requires the calculation of free energy differences between successive intermediate states generated using Eq. 6, which are later summed to provide $\Delta F_{0 \rightarrow 1}$ (SI 1). It is reported to

be less sensitive than other methods to the overlap deficiencies that might occur between two energy distributions obtained from two successive MD trajectories [47].

3 Application to the MD of Ubiquitin

Ubiquitin is a reference protein system that has already been studied by MD simulations [48-51]. It involves 76 AA residues and its secondary structure is characterized by a β -sheet made of five strands as well as two α -helices formed by residues 23 to 34 and 56 to 59. In the present work, molecular simulation conditions were kept as close as possible of those proposed by Showalter and Brüschweiler in their work about the Amber99SB FF [49]. MD trajectories of the system were run using the GROMACS 4.5.5 program package [35,36] with the Amber99SB FF [49] under particle mesh Ewald periodic boundary conditions. Long-range dispersion corrections to energy and pressure were applied. The initial configurations were retrieved from the Protein Data Base (PDB: 1UBQ) and solvated, when required, using TIP4P-Ew (a four-site model) [52] or SPC (a three-site model) [53] water molecules so as protein atoms lie at least at 1.2 nm from the cubic box walls. The His residue of Ubiquitin is in its His ϵ state, thus leading to a total protein charge of 0 [e⁻]. The systems were first approximately optimized to eliminate large forces and then heated to 50 K through a 10 ps canonical (NVT) MD, with a time step of 2 fs and LINCS constraints acting on bonds involving H atoms. The trajectory was followed by two successive 20 ps heating stages, at 150 K and at the final temperature, *i.e.*, 300, 277 or 250 K, under the same conditions. Next, each system was equilibrated during 50 ps in the NPT ensemble to relax the solvent molecules. Finally, a 20 ns MD simulation was performed in the NPT ensemble, for explicitly solvated systems. The ‘V-Rescale’ and ‘Parrinello-Rahman’ algorithms were selected to perform NVT and NPT simulations, respectively. Free energy differences between the reduced and all-atom models were evaluated during these 20 ns trajectories. An extra production run of 20 ns was performed for the evaluation of energetic, structural, and dynamical properties of the systems. Trajectory data were saved every 2 ps. A description of the systems under study is presented in Table 1. The total number of point charges to be considered for the protein is reduced by a factor of 4.3 and 12.3 for the CD- and PASA-based models, respectively. Depending upon the implementation, the number of non-atomic charges of Ubiquitin is largely variable. For instance, there are only two of such point charges in model mCDa, which originate from the Phe residues of Ubiquitin (Figure 1), while

the largest number is obtained for mPASA, with 99 non-atomic charges. There are, in each simulation, approximately 10350 water molecules that are not coarse-grained.

To evaluate the gain in calculation time, CPU times were compared for the vacuum MD simulations (Table 1). Values are reported for 10^7 MD steps carried out on two 2.66 GHz processors for the all-atom and CD-based models, and two 2.20 GHz processors for the PASA-based model. A gain of about 25 % in calculation time is observed. Keeping a significant number of all-atom contributions to the FF, *i.e.*, bonded and Lennard-Jones (LJ) terms, prevents a larger gain in the calculation time. For solvated systems, it is clear that a real gain in simulation time will be possible if coarse-graining occurs at the solvent level.

3.1 Protein structure

In our previous paper [34], the study of molecular electrostatic maps showed that at short range, *e.g.*, in the case of intramolecular interactions, the point charge models are expected to affect the dynamical behaviour of the molecules. As seen from the analysis of the MD trajectories of Ubiquitin, changes in the secondary structure elements are indeed observed (Figure 2). The figure illustrates that the structure of the protein is characterized by a loss of the secondary structure elements *versus* the all-atom simulation results, both in water and in vacuum. Among all RPCMs, mCD seems to lead to the most important loss in regular secondary structure elements when the system is solvated. Indeed, as already shown in ref. [34], mCDa is more efficient in the modelling of some secondary structure elements *versus* model mCD. mCDa allows to preserve the α -helix in SPC water, while mPASA preserves β -sheet elements, again in SPC water. In vacuum, β -strand and α -helix moieties disappear with models mCDa and mPASA, respectively. Only model mCD allows to preserve almost all secondary structure elements in vacuum. As mCD appeared to lead to the least stable structures in water, additional MD simulations were carried out at lower temperatures, *i.e.*, 277 and 250 K, like in our previous study [34]. The last temperature value was not selected to reflect a physical state for water (it is below the freezing point of the solvent) but was chosen to locally probe the potential energy hyper-surface of the system. At such low temperatures, stabilization of the secondary structure occurs for the mCD model, which illustrates that the energy hypersurface minimum observed with the all-atom model is existing with the mCD model, but energy barriers are lowered [34]. The existence of that minimum may be due to the conserved all-atom bonded and LJ energy terms in the FF. At 277 K, TIP4P-Ew water seems more structuring than SPC and more secondary structure

elements of 1UBQ are stabilized while, at 250 K, almost all secondary structure elements are present regardless of the water FF. Snapshots of the protein structure taken at the end of the last 20 ns MD trajectories (Figure 3) confirm a better secondary structure conservation for the mCD model in vacuum, at all studied temperatures, with RMSD values of 3.23, 2.52, and 2.05 Å, and in water at 250 K. RMSD values that are reported in Figure 3 were calculated versus the C α atoms of the PDB structure using the program VMD [54]. For the solvated systems, C α -C α distance maps established from the final snapshots are displayed in SI 2. Besides the all-atom calculations, only models mCDa and mPASA let a helix-like structure appear for residues 23 to 28, as well as β -strand contact regions, between strands 10 to 17, 1 to 7, 64 to 72, 40 to 45, and 48 to 50, for that particular MD frame. For mCD, it appears that the fold of the first half of the AA sequence only is preserved, except at low temperatures where the fold is recovered.

Other structural analyses of the MD trajectories obtained for the models dealt with the gyration radius r_G of the protein, which appears to be affected by the protein environment (Table 2). Nevertheless, a precise effect of the model on r_G is not clearly established. In water, r_G presents various trends. Stability is observed mainly with SPC water, except for mCD where the standard deviation is equal to 0.041 nm, while with TIP4P-Ew, both CD-based models, *i.e.*, mCD and mCDa, present slightly increasing r_G values, characterized by larger deviation values, 0.016 and 0.014 nm. Except for mPASA, all r_G values obtained in vacuum are smaller than in water, by about 5 % for the all-atom model, but up to 19 % for the mCD model. This was already reported in our last paper regarding MD simulations of the Ubiquitin-UIM-1 complex [34], where we showed that, in vacuum, the RPCMs led to a high protein compactness together with a lower atom mobility. The mPASA representation level seems to be the least affected by the presence of the solvent, in other words, r_G is similar, regardless of the environment. Such a lack of structuring effect depending on the medium was also illustrated in Figure 2.

H-bonds were determined based on cut-off values of 30° and 0.35 nm for the angle Hydrogen-Donor-Acceptor and the distance Donor-Acceptor, respectively. From values reported in Table 3, one first notices that the contraction of the protein in vacuum comes with a huge increase in the number of intra-molecular H-bonds, regardless of the model. For example, a mean value of 95.4 versus 55.5 H-bonds is observed for the all-atom model, in water and in vacuum, respectively. For the particular mCD model, one even observes that, at T = 250 K, the number of H-bonds in vacuum reaches the value obtained for the all-atom model, *i.e.*, a value close to 95 H-bonds. Second, even though the water model does not seem

to affect the number of intra-molecular H-bonds at the all-atom level, it is not the case for the RPCMs. Model mCD however appears to be the least affected, with values of 10.0 and 10.3 H-bonds under the TIP4P-Ew and SPC solvents, respectively. Third, all RPCMs lead to a drastic decrease in the number of such H-bonds *versus* the all-atom model, due to the cancelling of charges on most of the H atoms of the AAs. As further discussed, it comes with an increased number of protein-water H-bonds (except for mPASA) which confirms the strong influence of the solvent on the protein deconstruction in the case of the CD-based RPCMs.

H-bond distance and angle distributions of intra-molecular H-bonds formed with Ubiquitin are illustrated in Figure 4. From such distributions, one first notices that the angle distributions present a maximum at about 12.5 and 15.0° for the all-atom models only, in water and in vacuum, respectively. All RPCMs lead to a different angle distribution with no well-defined maxima, thus showing a loss in the orientational properties of H-bonds. Regarding the distances, mCD appears to best approach the distributions obtained for the all-atom model with, for example in TIP4P-Ew water, a maximum at 0.30 and 0.29 nm, respectively. mCD however leads to two slightly separated maxima when the SPC model is used, located at 0.29 and 0.31 nm. With TIP4P-Ew, mCDa also adopts two maxima, with minimal distances that are clearly shorter than in the all-atom distance distribution, *i.e.*, 2.39 *versus* 2.48 nm, while mCD has the same minimal value. In vacuum, there is a better agreement between the all-atom and the CD-based distance distributions. On the whole, even if point charges are not located on atoms in the mCD model, it does not appear to significantly affect H-bond distances.

A plot of C_{α} - C_{α} radial distribution functions (RDF) (Figure 5) shows a change in the structuring of the protein in the sense that, in water, a strong decrease of the very first peak, *i.e.*, around 0.6 nm, of the RDFs is observed for all RCPMs, especially mCD. SPC water emphasizes this effect. The use of CD-based models seems to involve a repulsion between the AAs, with RDF shortest distances that are now less probable when a solvent is used. It is even more marked in TIP4P-Ew RDFs, with only a shoulder at about 0.45 nm. All these observations are to be related to the enhanced loss in secondary structure elements already observed before (Figure 2). Contrarily, model mPASA appears to lead to the most similar RDFs *versus* the all-atom model, with however, a less structured long-range behaviour. In vacuum, the agreement between all RDFs is clearer.

The mCD model, with atomic charges located on virtual sites, thus involves stronger changes in the protein structure under the influence of explicit water molecules. When one studies the

charge occurrence distributions of Ubiquitin (Figure 6), one notices that mCD is characterized by the most positive values, just below $1.5 |e^-|$, as well as by a high content of the most negative charges, close to $-1 |e^-|$. mPasa leads to a high content of slightly negative charges, as in the all-atom model. mCDa is also characterized by two sets of more negative and more positive charges, but it seems that their location on atoms favours the preservation of secondary structure elements.

3.2 Protein hydration and H-bond dynamics

A study of the distance and angle distributions of the inter-molecular H-bonds formed between the solvated protein and water shows that CD-based models have a similar shape to the all-atom model while mPASA is the less consistent one (Figure 4). It is particularly clear when one studies the angle distributions wherein mPASA presents a maximum at about 27° rather than 9° , regardless of the water model. More surprisingly, for model mCD, a larger average number of main chain H-bonds, 245.5 *versus* 190.2, is obtained despite the absence of charges on the N and H atoms of the protein main chain (Table 4). It appeared to be due to the C=O groups that, with their different charge distribution *versus* the all-atom case, affect the formation of such a type of interaction, but also to the protein deconstruction [34], leading to a larger accessible surface for the formation of H-bonds with the solvent. The consequence of those changes in the number of H-bonds formed with the solvent is illustrated in Figure 7, where it is clearly seen that the very first hydration shell composed of water molecules interacting with the protein surface atoms, is levelled in the RPCM RDF curves. However, the minimal distance observed in the RDF is similar for all models, except for mPASA which appears to be more “repulsive” with values of about 0.17 *versus* 0.15 nm in the all-atom case. Such changes observed for the mPASA model are accompanied by huge decreases in the number of H-bonds, *i.e.*, 77.6 and 80.4 H-bonds for TIP4P-Ew and SPC, respectively, due to the absence of any dipole on the AA main chains (Table 4). One can finally mention that all H_w -protein RDFs are characterized by a shorter contact distance than their corresponding O_w -protein RDF, showing that H atoms are statistically closer to the protein structure.

The dynamics of protein-water H-bonds can be characterized through the so-called H-bond autocorrelation functions (SI 3):

$$C(t) = \frac{\langle h(0)h(t) \rangle}{\langle h \rangle} \quad (7)$$

where $h(t)$ is assigned a value of 1 or 0 if a particular pair of atoms is H-bonded or not. The approach that was applied to evaluate overall correlation times τ associated with $C(t)$, is:

$$\tau = \int_0^{\infty} C(t) dt \quad (8)$$

The H-bond autocorrelation functions were also fitted with a 3-exponential function:

$$C(t) = ae^{-t/\tau_1} + be^{-t/\tau_2} + ce^{-t/\tau_3} \quad (9)$$

with parameters listed in Table 5. All fits led to correlation coefficients R between 0.997 and 0.999. Visually, the mPASA function has the most similar behaviour to the all-atom one, while mCD and mCDa functions are characterized by a slower decrease (SI 3). This is consistent with the larger overall τ values reported in Table 5. For example, one gets values of 185.54 and 154.04 ps for models mCD and mCDa, respectively, to be compared with $\tau = 77.08$ ps for the all-atom model. Lowering the temperature has the expected effect to make relaxation slower, and thus τ larger. SPC and TIP4P-Ew have an opposite effect on τ , increasing its value for mCDA or mCD when TIP4P-Ew or SPC are applied, respectively. A detailed study of parameters of Eq. 9 shows that the influence of the water FF becomes more important at larger times, which may be due to an effect of the protein conformational relaxation. Such an influence is also visible for mCDa. The effect on mPASA is seen through the well-marked increase of parameter a versus b . When T decreases, the third component of $C(t)$, that is associated with the largest correlation time, is increased versus a and b . For example, for mCD with TIP4P-Ew at 300, 277, and 250 K, one gets $c = 0.188$, 0.218, and 0.401, respectively, while τ_3 has no precise trend. This is also observed with SPC. For the fastest contribution, mPASA differs the most versus the all-atom case, with a higher proportion of the faster component $a = 0.577$, and a lower value of the corresponding correlation time $\tau_1 = 1.55$ ps. With SPC, differences between mCD and mCDa are limited, while they are larger with TIP4P-Ew, especially for τ_1 .

3.3 Energetics and thermodynamics

MD-averaged energy values are not easily comparable between the different models as the protein conformations generated by the MD simulations differ. A first discussion is thus based on the comparison of potential energy values E of Ubiquitin calculated from all intra-molecular energy terms for the conformations generated through MD trajectories. Values were obtained using post-processing calculations applied to the TIP4P-Ew 20 ns MD trajectories of the RPCMs (SI 4). The correlation coefficient values obtained for $E_{\text{all-atom}}$ versus E_{mCD} , $E_{\text{all-atom}}$ versus E_{mCDa} , and $E_{\text{all-atom}}$ versus E_{mPASA} are equal to 0.914, 0.902, and 0.617, respectively. The values illustrate the better agreement with Ubiquitin all-atom potential energy values for the CD-based RPCMs, but still show that part of the conformational energies do not adopt the same trend as in the all-atom model.

A second discussion concerns the optimized, and thus similar, protein conformations (SI 5). Regarding intra-molecular energy contributions, the reduction of the point charge number leads to a decrease of bonding energy terms, *i.e.*, stretching, bending, dihedral, versus their corresponding all-atom contribution. For example, the stretching terms of the RPCMs are about 80 to 90 $\text{kJ}\cdot\text{mol}^{-1}$ versus 137.42 $\text{kJ}\cdot\text{mol}^{-1}$ for the all-atom model. This comes with an increase of the non-bonding Coulomb (Cb) and Lennard-Jones (LJ) terms. Cb-14 contribution, *i.e.*, Cb interactions involving two atoms separated by three chemical bonds, is absent in the mPASA model while it is systematically larger by about 3200 $\text{kJ}\cdot\text{mol}^{-1}$ for the mCD model, consistently with the charge distributions analysis reported earlier (Figure 6). This increase is also observed for the MD simulations, even at low temperatures. Thus, mCDa with a value of 12465.32 $\text{kJ}\cdot\text{mol}^{-1}$ is closer to the all-atom value, *i.e.*, 12463.77 $\text{kJ}\cdot\text{mol}^{-1}$, due to the preserved location of the charges on the atoms of the protein structure. The short-range LJ term (LJ-SR) of mCD appears to also be less stabilizing, regardless of the medium, except at lower temperatures. Indeed, for mCD, one notices an average of -1531.84 versus -2203.11 $\text{kJ}\cdot\text{mol}^{-1}$ for the TIP4P-Ew all-atom case. Similarly, the Cb-SR term is less stabilizing, whatever the medium is, and the mCD term is closer to the corresponding all-atom term, with -3944.84, -3841.82, and -6572.02 $\text{kJ}\cdot\text{mol}^{-1}$ for the optimized, TIP4P-Ew solvated, and vacuum structures. When solvation is considered, the TIP4P-Ew FF appears to provide intra-molecular energy values that are closer to the corresponding values for the optimized structures, except for mPASA.

Regarding inter-molecular protein-water interactions, the CD-based RPCMs emphasize the negative character of the Cb-SR term at all temperatures, while mPASA is largely less

stabilizing. For example, mean values of -8024.49, -12127.84, -11436.43, and -3423.91 kJ.mol⁻¹, are obtained for the TIP4P-Ew all-atom, mCD, mCDa, and mPASA models, respectively. The change in the protein conformation together with the change in the point charge model lead to varying LJ terms. In water, LJ-SR is stabilizing for mCD and mPASA *versus* the all-atom, with mean values of -708.75, -1026.21, and -506.14 kJ.mol⁻¹, respectively. The greater influence of the solvent for these two RPCMs involves a change of 186.54 kJ.mol⁻¹ for mCD and -60.56 kJ.mol⁻¹ for mCDa, while it is only 6.32 kJ.mol⁻¹ for the all-atom model. Again, a clear comparison is difficult to achieve because protein conformations differ.

Free energy differences between the all-atom and RPCMs were calculated using the two different approaches briefly described earlier. With the BAR method, 11 simulations of 10⁷ steps each were carried out for λ values between 0 and 1 with a step of 0.1, λ being kept constant at each simulation. The approach was applied only to systems in vacuum. With the slow growth approach, one simulation of 10⁷ steps was achieved, with λ varying from 0 to 1 with a step of 10⁻⁷, λ being changed at each simulation step. Values of $\Delta G_{all-atom \rightarrow RPCM}$ are reported in Table 6. It is seen that the slow growth approach provides values that are comparable to BAR ones, even if slightly larger, by about 2 % for the mCD model at 300 K to 7.5 and 10.9 % at 277 and 250 K, respectively. This can be due to the fact that the two states under study, the all-atom and the RPCM ones, differ only by their point charges. To understand the range of values obtained for the RPCMs, calculations were also achieved for rigid systems, *i.e.*, systems for which the conformational contribution to ΔG should be limited, allowing the evaluation of the electrostatic contribution to ΔG .

It is first observed that all ΔG values are positive, denoting a favoured all-atom point charge representation, even for rigid systems. Rigidity involves very different trends. In vacuum, ΔG is largely reduced *versus* the flexible case. For example, one obtains ΔG differences of (748.86 - 1658.86) kJ.mol⁻¹ for the mCD system. In the presence of TIP4P-Ew water, ΔG increases, with now a ΔG difference of (1685.07 - 1313.26) kJ.mol⁻¹. SPC seems to bring very little changes between the rigid and flexible systems. As ΔG values are higher for systems held fixed in TIP4P-Ew water, it is concluded that the deconstruction of the protein structure during the simulation has a favourable effect on the all-atom-to-RPCM transformation. Contrarily, in vacuum, flexibility is unfavourable to the transformation. It might be related to the better structure conservation of mCD in vacuum than in water

(Figures 2 and 3). As observed for previously studied properties, mPASA differs very much from the other RPCMs, in that the ΔG values are very high, around 7 to 9 10^3 kJ.mol⁻¹, and the model leads to a larger ΔG in water than in vacuum, contrarily to the other RPCMs. For example, the presence of water favours the alchemical all-atom-to-mCD transformation by 345.60 kJ.mol⁻¹, with ΔG equal to 1658.86 kJ.mol⁻¹ in vacuum and 1313.26 kJ.mol⁻¹ in TIP4P-Ew water. Contrarily to some previously studied structural properties, mCD now appears to be closer to the all-atom model, both for the rigid and flexible systems.

The study of rigid systems allows to evaluate the electrostatic contribution to ΔG , ΔG_e , and its intra- and inter-molecular components $\Delta G_{e-intra}$ and $\Delta G_{e-inter}$. For mCD, $\Delta G_{e-intra} = 748.86$ kJ.mol⁻¹, *i.e.*, ΔG calculated for the rigid system in vacuum, while $\Delta G_{e-inter}$ is assumed to be the difference between the corresponding ΔG value for the solvated system minus the previous value, *i.e.*, $1685.07 - 748.86 = 936.21$ kJ.mol⁻¹. Following such a reasoning, SPC has thus a smaller effect on the $\Delta G_{e-inter}$ values than TIP4P-Ew.

In literature, values for the electrostatic contribution to the solvation free energy of Ubiquitin, ΔG_{e-solv} , are reported [55,56]. Depending upon the method used to evaluate ΔG_{e-solv} , values of -997.02, -1049.17, -1085.22 kcal.mol⁻¹ [55], -1446 and -1459 kcal.mol⁻¹ [56] are given. If one considers the all-atom value obtained with the TIP3P and Amber94 potentials, *i.e.*, -1085.22 kcal.mol⁻¹, a thermodynamical cycle (Figure 8) provides a value of -3604.36 kJ.mol⁻¹ for the hydration free energy of the mCD protein, a value lower by 936.17 kJ.mol⁻¹ than the all-atom value (Table 6). Such differences cannot be fully explained by the charge distributions (Figure 6), as model mPASA presents a charge distribution that is similar to the all-atom one.

3.4 Water dynamics

To study the effect of the point charge model on water dynamics, solvent molecules were classified in two sets, within 0.35 nm from the protein atoms, and between 0.35 and 1.40 nm. As water molecules can migrate from one region to another during the simulation, the following strategy was employed. First, at each 1000th frame (every 1 ns), water molecules included in a layer of 0.35 nm from the protein surface atoms were detected. Then, the number of molecules staying in the given layer, for a given period of time, were averaged, considering each frame as a time origin (Table 7). From Table 7, it is first seen that the mean number of water molecules interacting closely with the protein surface is higher for the CD-based RPCMs than for the all-atom model. For example, one observes averages of 546.1

molecules for the mCD model rather than 358.5 for the all-atom one. Among those values, about 1.7 and 2.6 % are still present in the layer after 5 ns, respectively. It thus confirms that water dynamics is slowed down when one uses a CD-based RPCM rather than the mPASA model that shows a faster water exchange with time. In that later case, fewer molecules, an average of 329.3, are actually observed in the layer. Conclusions are similar when SPC is used, with however a slightly faster exchange as discussed hereafter, notably due to a larger water self-diffusion coefficient, as illustrated later. Contrarily, a slower exchange is observed, **as expected**, at lower temperatures for the mCD model.

Persistence times, characterizing a fast and a slower regime (Table 8), were obtained by fitting a two-exponential function to the normalized mean number of water molecules, $N(t)$:

$$N(t) = ae^{-t/\tau_1} + be^{-t/\tau_2} \quad (10)$$

All fits led to a correlation coefficient R of 0.9997 or higher. It is assumed that the shortest persistence times, τ_1 , are associated with molecules that are at the boundary of the water layer, and are thus exchanging faster with the bulk, while the largest persistence time, τ_2 , characterizes water molecules that are more strongly bound to the protein surface atoms through H-bonds. The major component, depicted by the larger parameter a , is assigned the shortest time constant τ_1 . In that framework, it is shown that CD-based RCPMs involve a slower dynamics (larger τ_1 values) than the all-atom model. For example, mCD and all-atom models give values of 0.423 and 0.337 ps, respectively, with the TIP4P-Ew potential. The mPASA τ_1 values are similar to the all-atom values, both for TIP4P-Ew and SPC solvent. mCD again appears as more easily influenced by the water FF than the other RCPMs, mCDa and mPASA, especially regarding τ_2 . A second component in the molecular behaviour is characterized by the smaller parameters b and by greater persistence times (slower dynamics). The values τ_2 however comes with the largest relative errors in the fitting program (up to 50 %) and are thus less reliable.

The hereabove observations are consistent with a study of self-diffusion coefficients of water in the layer. Again, to be certain to study molecules that are within the layer, a short time range was considered, from 10 to 11 ns. This choice allowed to consider a relatively large number of molecules. The self-diffusion coefficients D were obtained through a fit of the MSD function between 100 and 400 ps. Results are presented in Table 9.

Dastidar and Mukhopadhyay also reported dynamical results for water as a function of their distance *versus* the Ubiquitin surface [48]. Values obtained for the all-atom model with the TIP4P-Ew water potential are consistent with their observations, *i.e.*, a decrease in D as water gets closer to the protein surface, with a similar magnitude. It is also seen, from our results, that all RPCMs but mPASA involve a decrease in D *versus* the all-atom model with, for example, $D = 0.64 \times 10^{-5}$ and $1.28 \times 10^{-5} \text{ cm}^2 \cdot \text{s}^{-1}$ for the mCD and all-atom models, respectively. This is consistent with the longer persistence times (Table 8), longer H-bond correlation times (Table 5), and more attractive Cb and LJ short range interactions (SI 5) discussed previously.

4 Conclusions and perspectives

Two reduced point charge models were considered for Molecular Dynamics (MD) simulations of Ubiquitin. The first model, based on charges located at critical points (CP) of smoothed amino acid (AA) charge density (CD) distribution functions calculated from Amber99 atomic values, involves two point charges on the main chain of each AA, precisely located on atoms C and O, and up to six charges for the side chain. The second model, built by assigning charges to the maxima of AA smoothed promolecular electron density distribution functions, considers one point charge on the main chain and no more than two charges on the side chain.

For the first model, two different implementations were considered. In a first stage, the model is applied as is by considering charges as virtual sites in the system (model mCD). Second, rather than being located away from atom positions, most of the charges are set at selected atom positions. Their values are recalculated accordingly (model mCDa). For the second model, only the first kind of implementation was considered (model mPASA).

MD simulations were carried out using the program GROMACS with the Amber99SB force field (FF), in water and in vacuum. Two water FFs were considered, TIP4P-Ew and SPC. The selected temperature was 300 K, except for model mCD for which two lower temperature values, 277, 250, and 150 K, were also considered.

Energetic, structural, and dynamical information were retrieved from the analysis of the MD trajectories of the reduced point charge models (RPCMs) and discussed *versus* the all-atom model and available literature data. An emphasis was put on the secondary structure elements of the protein, the energetics and free energy difference between RPCMs and the all-atom model, and the characterization of H-bonds within the protein and with the solvent.

On a structural point of view, one observed a progressive loss in the secondary structure of the proteins when RPCMs are used at room temperature. However, model mCD applied at low temperatures with the TIP4P-Ew FF, or in vacuum, allows to preserve the secondary structure elements of Ubiquitin. It is explained by a free energy difference in favour of a spontaneous transformation from the mCD to the all-atom state in vacuum, while the inverse transformation is favoured in water. At 300 K, model mCDa better preserves some secondary elements, due to a better description of the short range 1-4 Coulomb and Lennard-Jones energy terms. Model mPASA led to the largest differences *versus* the all-atom model in terms of energetic, structural, and dynamical properties of the system, as also illustrated in [ref. \[34\]](#). First, the implementation of the mPASA model is such as no Cb-14 contributions to the potential energy are involved. Second, the number of point charges is too low to allow a first hydration shell as in the all-atom and in the other RPCMs. Geometrical parameters, that are associated with the existing H-bonds, adopt larger distance and angle values. Additionally, these H-bonds show a faster dynamics. The structure of the protein is particularly deconstructed in vacuum, contrarily to the trends followed by the other models. Even in water, Ubiquitin undergoes the more important contraction effect with the smallest gyration radius, while the other RPCMs systematically lead to an increase of the radius as shown in our previous paper [34].

If the CD-based RPCMs do not favour the formation of a first hydration shell as clearly as the all-atom model does, they however allow the formation of solute-solvent H-bonds with geometrical properties similar to the all-atom case. Intra-protein H-bonds are differently described with no well-defined angle distributions. On the whole, the change in the water FF has little effect on the all-atom protein model, but it has the largest effect on the mCD structure and thermodynamics. Indeed, TIP4P-Ew is best to model the protein structure and is more structuring at low temperature, possibly due to low self-diffusion coefficients D . SPC provides, on the average, a larger repulsion between the AAs, as corresponding C_{α} - C_{α} radial distribution functions (RDFs) show less probable short separation distances. It also accelerate the protein-water H-bond dynamics.

MD simulations carried out with model mCD at temperatures below 300 K led to the conclusion that this particular point charge model is able to provide results that are essentially similar to the all-atom model. One finds there a clue to conclude that, with RPCMs, energy barriers of the potential well are lowered, conformations can be perturbed more easily, but the location of that potential well on the energy hyper-surface is similar [34]. Model mCD is also the most similar, in terms of free energy, to the all-atom one.

In conclusions, all aspects of MD analysis, *i.e.*, energetics, dynamics, and structural criteria should be considered together to assess a RPCM, as they may give various trends. As already stated in [ref.](#) [34], Cb-14 term is important to preserve the protein structure. These terms are better reproduced when charges are located on atoms. Cb-SR seems also to be a characteristic that differentiates all-atom and RPCMs. As a perspective, a revision of the virtual site definition might be a path to follow in order to modify the Cb-14 energy terms to allow better RPCM MD trajectories. A way to revise such interactions is to modify the reference atoms selected to define the virtual sites. Indeed, within the GROMACS implementation, the forces acting on the virtual sites are redistributed among the reference atoms. Another aspect is to revisit the calculation of the point charge values.

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Figure captions

Figure 1 Description of the three reduced point charge models (black spheres) of selected AA residues in the framework of the Amber99 FF.

Figure 2 Secondary structure of Ubiquitin observed during the last 20 ns Amber99SB-based MD trajectories at 300 K, obtained using the all-atom, mCD, mCDa, and mPASA models. Secondary structure elements are colour-coded as follows: Coil (white), α -helix (blue), π helix (purple), 3_{10} helix (grey), β -sheet (red), β -bridge (black), bend (green), turn (yellow). (For an interpretation of the references to colour, please refer to the online version of the paper).

Figure 3 Final snapshots of Ubiquitin obtained from the last 20 ns Amber99SB-based MD trajectories at 300 K, generated using the all-atom, mCD, mCDa, and mPASA models. Secondary structure elements are colour-coded as follows: Coil (white), α -helix (blue), π helix (purple), 3_{10} helix (grey), β -sheet (red), β -bridge (black), bend (green), turn (yellow). Figures were generated using VMD [54]. RMSD values are calculated *versus* the C_α coordinates of the PDB structure. (For an interpretation of the references to colour, please refer to the online version of the paper).

Figure 4 Distance and angle distribution functions of intra- and inter-molecular H-bonds formed with Ubiquitin calculated from the last 10 ns of the Amber99SB-based MD trajectories at 300 K, obtained using the all-atom, mCD, mCDa, and mPASA models under various solvation conditions.

Figure 5 RDF of C_α - C_α atom pairs of Ubiquitin calculated from the last 10 ns of the Amber99SB-based MD trajectories at 300 K, obtained using the all-atom, mCD, mCDa, and mPASA models under various solvation conditions.

Figure 6 Occurrence distributions of atomic charges, $p(q)$, of Ubiquitin for the all-atom, mCD, mCDa, and mPASA models.

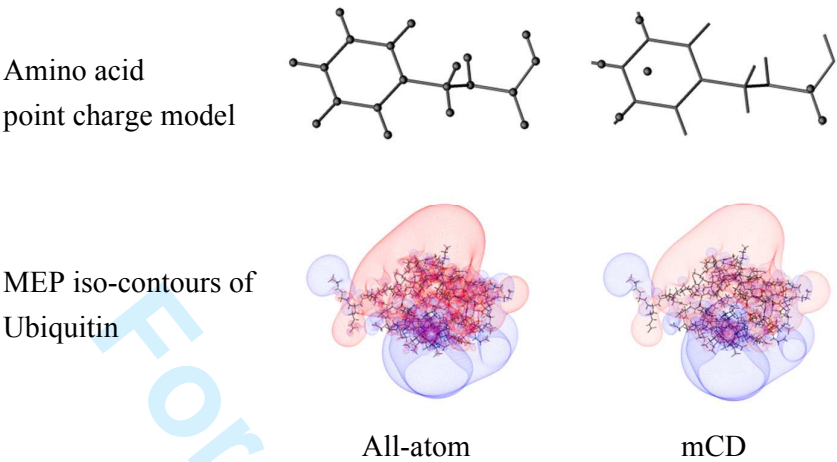
Figure 7 RDF of water oxygen- and water hydrogen-protein atom pairs of solvated Ubiquitin calculated from the last 10 ns of the Amber99SB-based MD trajectories at 300 K, obtained

using the all-atom, mCD, mCDa, and mPASA models with TIP4P-Ew and SPC water FFs. All protein atoms are included in the calculations.

Figure 8 Thermodynamical cycle used for the evaluation of the electrostatic contribution to the free energy of solvation of Ubiquitin, ΔG_{e-solv} , under the TIP4P-Ew potential, at 300 K.

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Graphical abstract



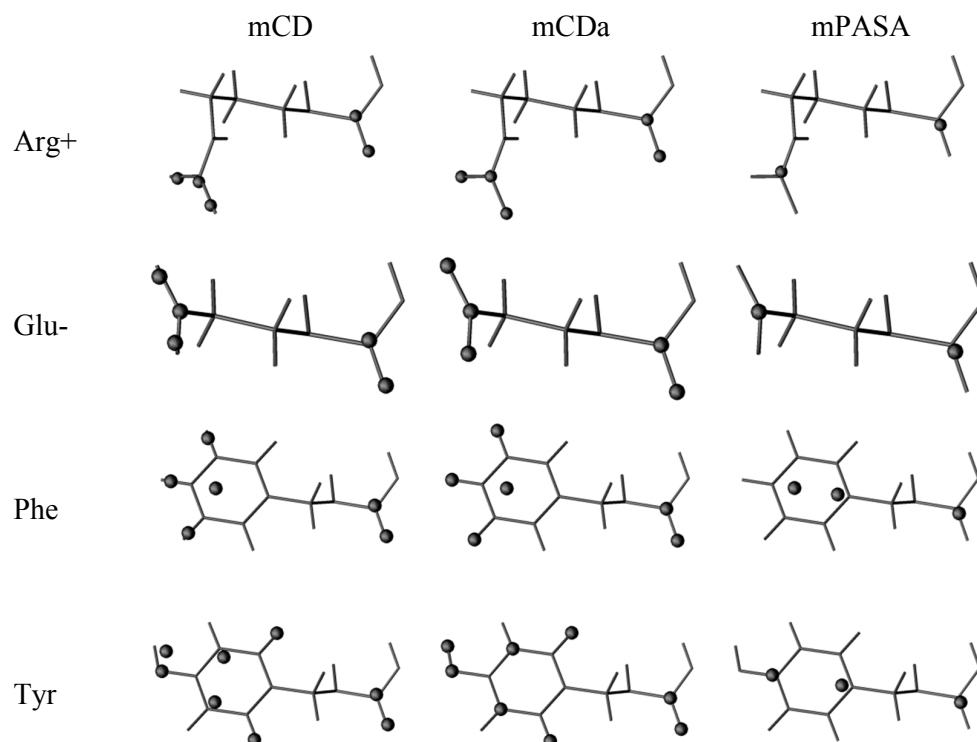


Figure 1.

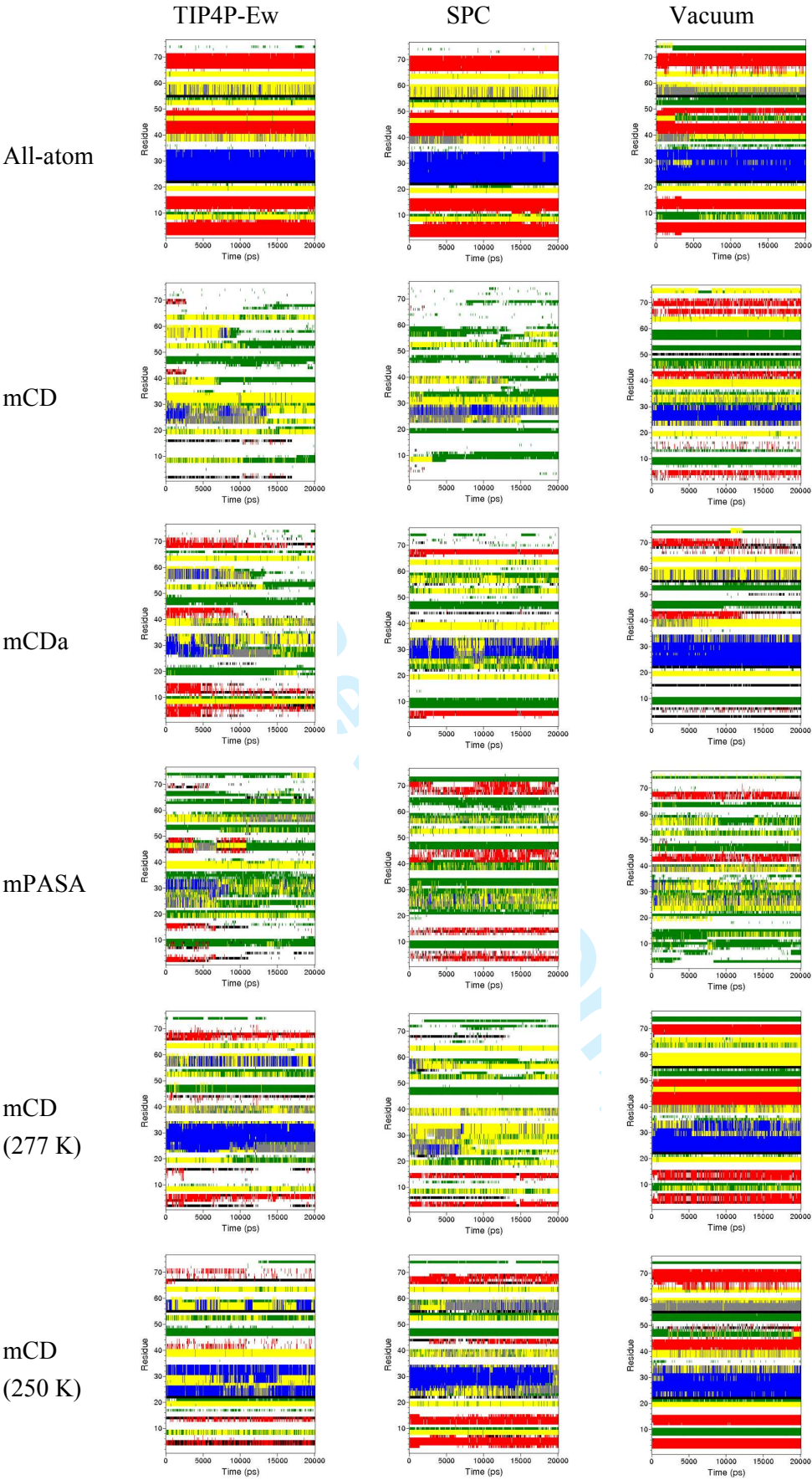


Figure 2.

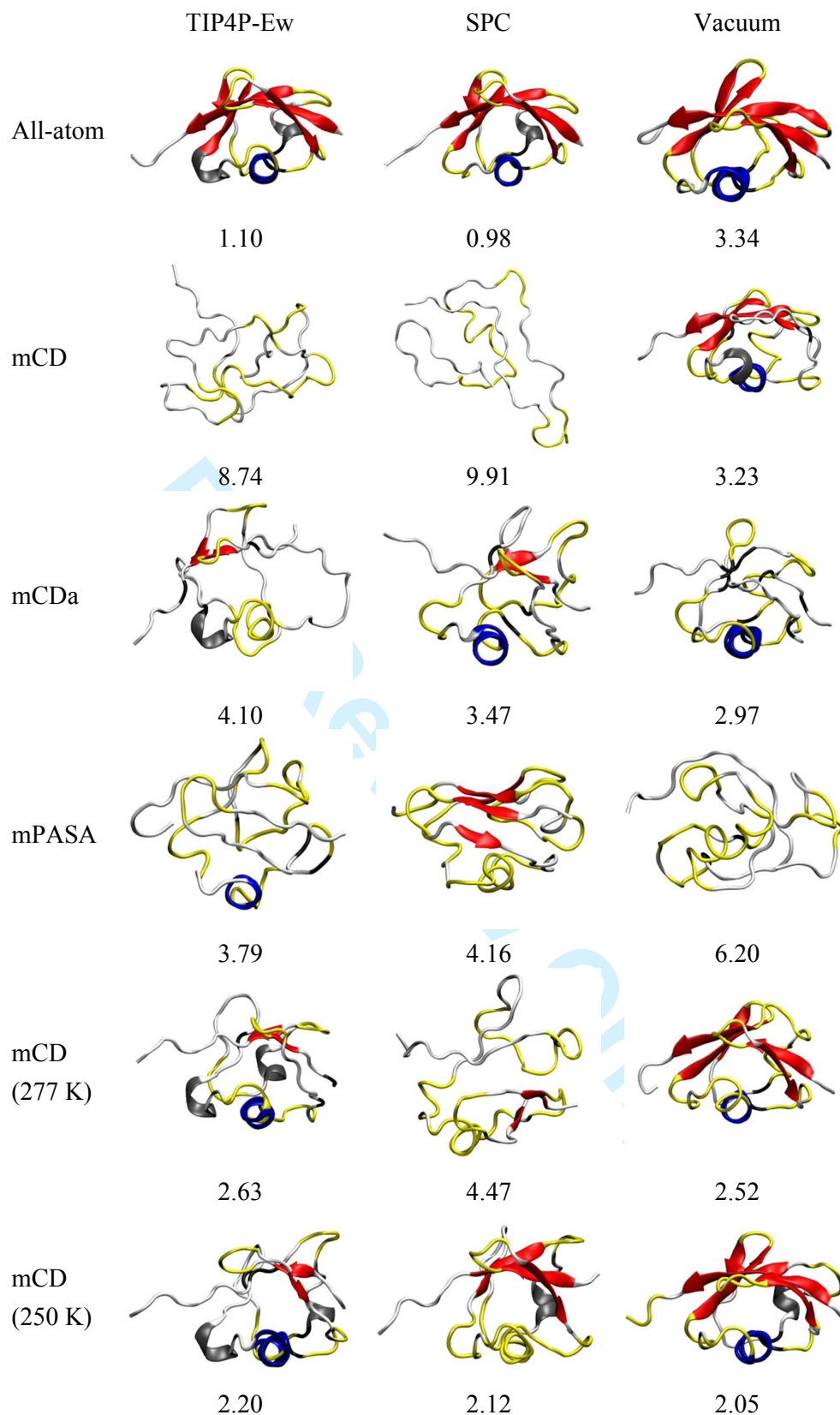


Figure 3.

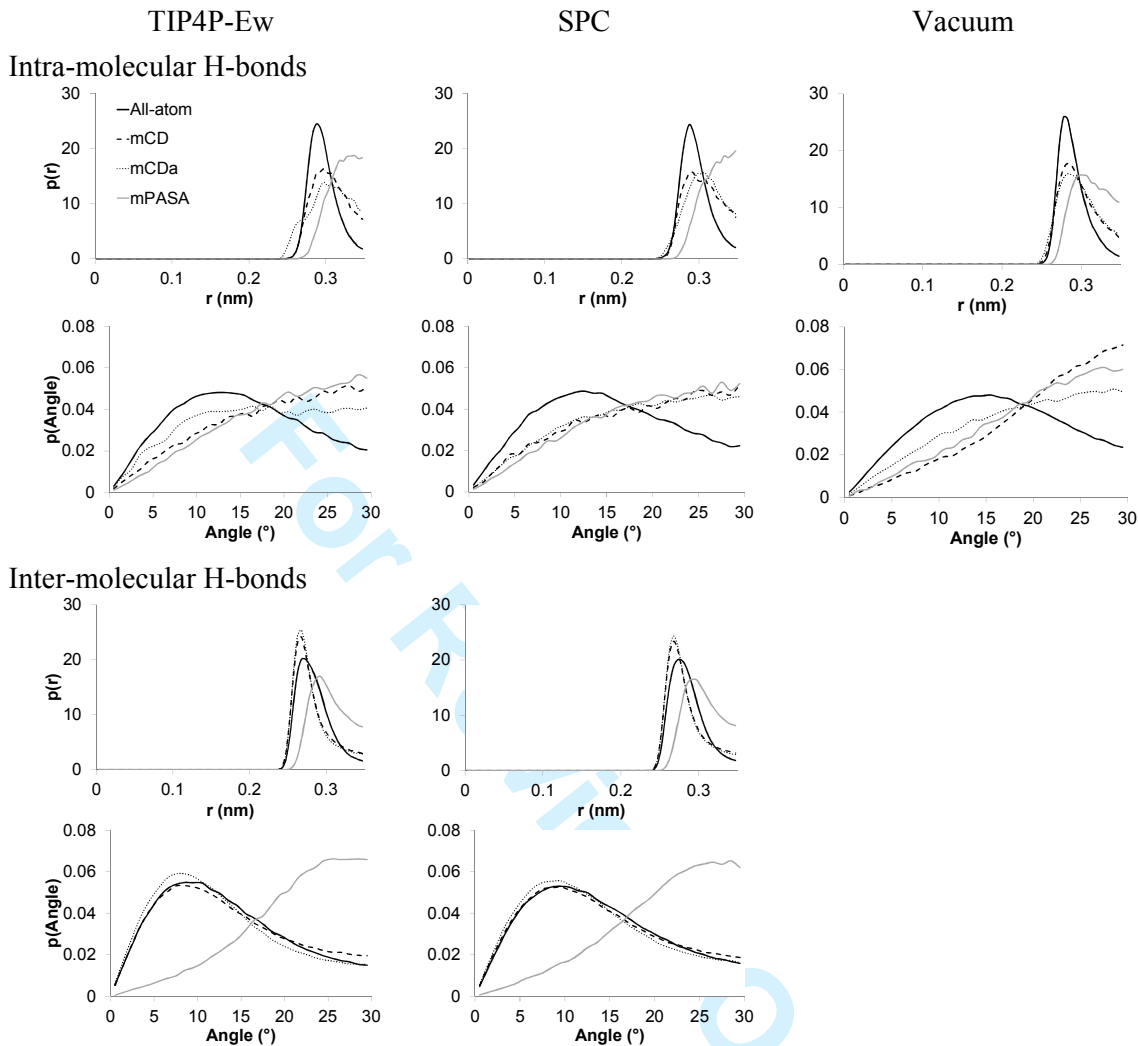


Figure 4.

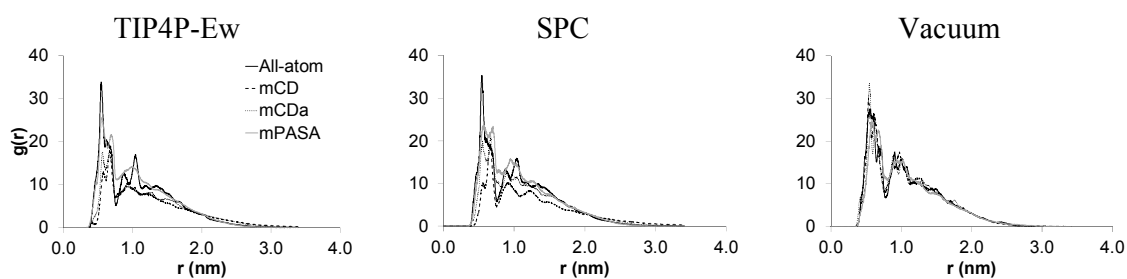


Figure 5.

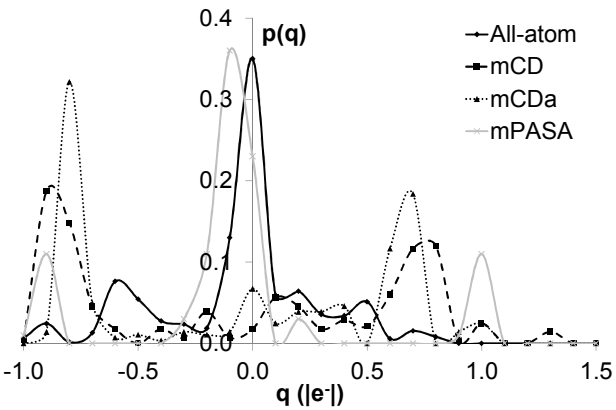


Figure 6.

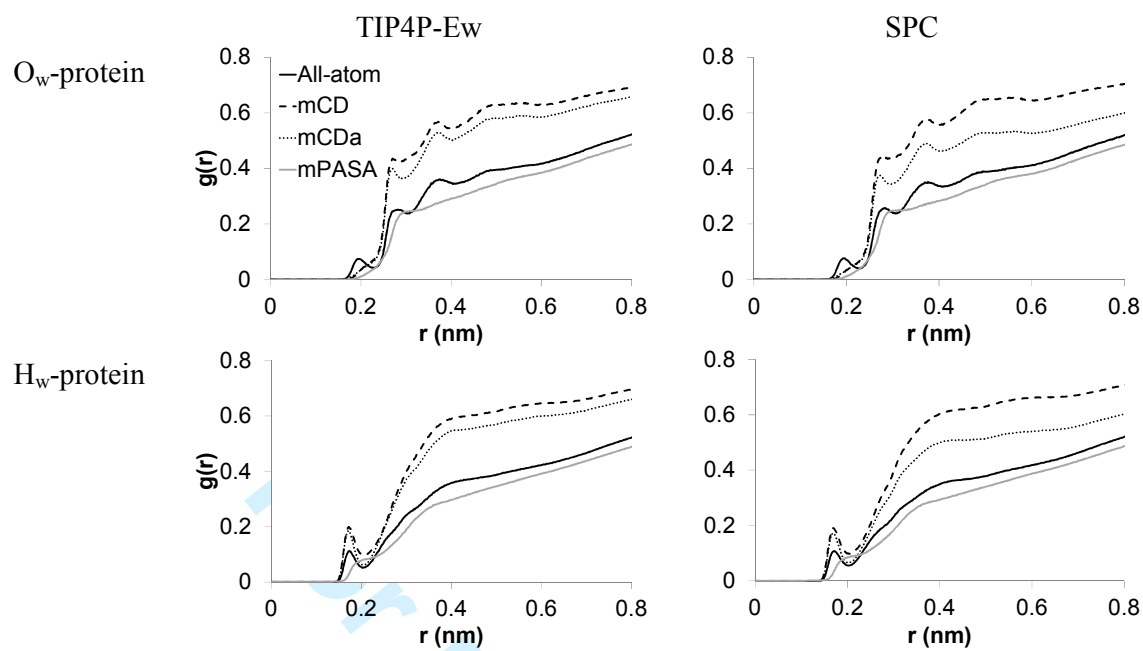


Figure 7.

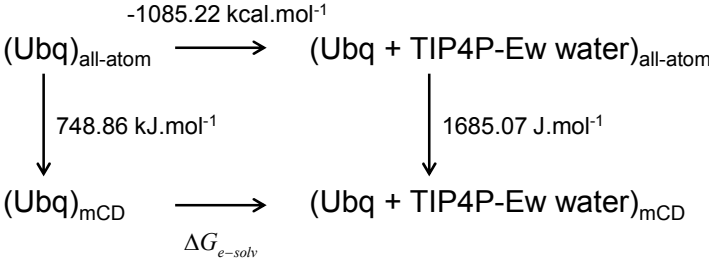


Figure 8.

Table 1 Description of the point charge models used for the Amber99SB-based MD simulations of Ubiquitin.

	No. of water molecules		No. of point charges associated with Ubiquitin	No. of non-atomic point charges	Box size (nm) (from final snapshot)	CPU time for vacuum simulations (h:m)
	TIP4P-Ew	SPC				
All-atom	10369	10346	1231	0	6.874	21:23
mCD	10366	10346	283	84	6.889	16:41
mCDa	10368	10346	283	2	6.890	15:55
mPASA	10368	10349	100	99	6.893	15:52

Table 2 Mean gyration radii r_G (nm), and their standard deviation, of Ubiquitin, obtained from the analysis of the last 10 ns of the Amber99SB-based MD trajectories at 300 K.

	Solvated		Vacuum
	TIP4P-Ew	SPC	
All-atom	1.193 ± 0.007	1.188 ± 0.007	1.127 ± 0.003
mCD	1.399 ± 0.016	1.492 ± 0.041	1.138 ± 0.004
mCDa	1.335 ± 0.014	1.248 ± 0.010	1.120 ± 0.005
mPASA	1.148 ± 0.009	1.169 ± 0.011	1.157 ± 0.007
mCD (277 K)	1.265 ± 0.009	1.300 ± 0.015	1.133 ± 0.004
mCD (250 K)	1.245 ± 0.007	1.237 ± 0.006	1.140 ± 0.004

Table 3 Average numbers of intra-molecular H-bonds, and their standard deviation, occurring in Ubiquitin, obtained from the analysis of the last 10 ns of the Amber99SB-based MD trajectories at 300 K.

	Solvated		Vacuum
	TIP4P-Ew	SPC	
All-atom	55.5 ± 3.1	55.5 ± 3.3	95.4 ± 3.2
mCD	10.0 ± 3.0	10.3 ± 2.4	38.0 ± 4.2
mCDa	12.6 ± 2.7	18.0 ± 3.1	41.5 ± 4.0
mPASA	11.4 ± 3.0	12.9 ± 3.3	20.0 ± 3.4
mCD (277 K)	18.3 ± 3.3	13.9 ± 2.7	87.5 ± 3.9
mCD (250 K)	17.9 ± 2.7	20.8 ± 3.1	93.4 ± 3.6

Table 4 Average numbers of H-bonds, and their standard deviation, occurring between Ubiquitin and water, obtained from the analysis of the last 10 ns of the Amber99SB-based MD trajectories at 300 K.

	TIP4P-Ew	SPC
All-atom	190.2 ± 6.6	190.3 ± 6.7
mCD	245.5 ± 10.1	240.0 ± 9.4
mCDa	230.9 ± 7.7	220.5 ± 7.3
mPASA	77.6 ± 6.4	80.4 ± 6.9
mCD (277 K)	230.3 ± 7.9	250.2 ± 8.7
mCD (250 K)	211.3 ± 6.9	220.6 ± 6.8

Table 5 Fitting parameters for the Ubiquitin-water H-bond autocorrelation functions (Eq. 9) obtained from the analysis of the last 10 ns Amber99SB-based MD trajectories at 300 K.

	all-atom	mCD	mCDa	mPASA	mCD (277 K)	mCD (250 K)
TIP4P-Ew						
a	0.423	0.385	0.414	0.577	0.364	0.272
b	0.481	0.418	0.408	0.310	0.408	0.325
c	0.087	0.188	0.158	0.113	0.218	0.401
τ_1 (ps)	4.92	4.34	6.15	1.55	4.40	3.03
τ_2 (ps)	53.62	83.88	82.02	59.90	106.17	121.74
τ_3 (ps)	554.45	821.49	765.84	296.97	909.43	870.58
Eq. 11 (ps)	77.08	185.54	154.04	54.10	232.77	373.34
SPC						
a	0.517	0.433	0.458	0.621	0.385	0.347
b	0.424	0.428	0.411	0.350	0.387	0.395
c	0.051	0.133	0.123	0.028	0.221	0.252
τ_1 (ps)	4.41	3.94	4.45	1.59	4.13	3.75
τ_2 (ps)	43.53	66.06	63.87	35.55	80.35	100.14
τ_3 (ps)	506.06	533.95	894.91	388.64	893.61	957.57
Eq. 11 (ps)	48.95	102.43	133.00	26.02	220.73	268.11

Table 6 $\Delta G_{all-atom \rightarrow RPCM}$ (kJ.mol⁻¹) calculated using the slow growth approach (BAR results are in parentheses) from Amber99SB-based 20 ns MD trajectories of Ubiquitin at 300 K. ΔG_{e-solv} and $\Delta\Delta G_{e-solv}$ are calculated for the TIP4P-Ew potential. ΔG_{e-solv} and $\Delta\Delta G_{e-solv}$ are based on a all-atom ΔG_{e-solv} of -1085.22 kcal.mol⁻¹ [55].

	Vacuum	Solvated		ΔG_{e-solv} (Figure 7)	$\Delta\Delta G_{e-solv}$
		TIP4P-Ew	SPC		
Flexible					
mCD	1658.86 (1631.82 ± 6.73)	1313.26	1367.71		
mCDa	1946.28 (1877.99 ± 5.59)	1491.10	1559.34		
mPASA	8215.53 (7756.48 ± 4.56)	8842.43	8751.73		
mCD (277 K)	1742.48 (1620.56 ± 4.47)	1352.01	1384.35		
mCD (250 K)	1816.12 (1637.22 ± 10.02)	1365.80	1357.78		
Rigid					
mCD	748.86	1685.07	1358.18	-3604.36	-936.17
mCDa	761.03	1703.65	1505.82	-3597.94	-942.62
mPASA	7346.06	9162.80	8731.14	-2723.82	-1816.74

Table 7 Mean number of water molecules staying in a layer of thickness 0.35 nm from the Ubiquitin surface for a given period of time. Values were obtained from the last 10 ns of Amber99SB-based MD trajectories at 300 K with the TIP4P-Ew and SPC water potentials. Values in parentheses are the ratios (%) of persistent molecules.

t (ns)	All-atom	mCD	mCDa	mPASA	mCD (277 K)	mCD (250 K)
TIP4P-Ew						
0	358.5	546.1	505.7	329.3	482.7	450.7
1	20.8	75.7	69.2	14.7	104.2	156.5
2	2.5	29.1	26.8	1.2	53.0	78.9
3	1.2	16.4	16.5	1.0	40.9	50.4
4	1.0	11.4	10.6	0	35.7	37.0
5	1.0 (2.6)	9.4 (1.7)	7.0 (1.4)	0	32.2 (6.7)	30.5 (6.8)
6	0	7.8	5.0	0	29.6	26.0
7	0	6.6	2.8	0	27.3	22.3
8	0	5.7	2.0	0	25.3	19.3
9	0	5.0	2.0	0	24.0	18.0
10	0	5.0	0	0	22.0	16.0
SPC						
0	349.2	554.9	452.2	320.5	514.5	452.6
1	13.7	51.5	38.2	11.9	80.4	93.5
2	1.7	11.4	13.7	1.0	35.4	44.6
3	1.2	4.8	9.6	0	23.8	33.4
4	1.0	2.6	7.4	0	18.7	29.3
5	1.0 (0.3)	2.0 (0.4)	6.0 (1.3)	0	15.3 (3.0)	26.5 (5.9)
6	1.0	1.7	5.4	0	12.6	24.2
7	1.0	1.0	5.0	0	10.5	22.3
8	0	0	4.7	0	7.7	21.7
9	0	0	4.5	0	6.5	21.0
10	0	0	4.0	0	5.0	20.0

Table 8 Fitting parameters for the normalized mean number of water molecules $N(t)$ located in a layer of 0.35 nm, at 300 K, from the protein surface as a function of time (Eq. 10).

	all-atom	mCD	mCDa	mPASA	mCD (277 K)	mCD (250 K)
TIP4P-Ew						
a	0.990	0.934	0.960	0.994	0.891	0.846
b	0.010	0.066	0.043	0.006	0.108	0.154
τ_1 (ps)	0.337	0.423	0.445	0.314	0.494	0.740
τ_2 (ps)	2.548	4.040	6.725	2.873	11.01	6.312
SPC						
a	0.994	0.956	0.966	0.996	0.907	0.910
b	0.006	0.044	0.034	0.004	0.093	0.090
τ_1 (ps)	0.296	0.376	0.351	0.298	0.417	0.503
τ_2 (ps)	4.879	1.907	6.423	0.764	4.42	12.659

Table 9 Self-diffusion coefficients ($10^{-5} \text{ cm}^2 \cdot \text{s}^{-1}$) of water, and their standard deviation, obtained from the last 10 ns of Amber99SB-based MD trajectories at 300 K with the TIP4P-Ew and SPC water potentials. Values were calculated over a 1 ns fragment of the trajectory and over the number of molecules given in parentheses.

	all-atom	mCD	mCDa	mPASA	mCD (277 K)	mCD (250 K)
TIP4P-Ew						
layer 0-0.35 nm	1.28 ± 0.11 (22)	0.64 ± 0.13 (74)	0.55 ± 0.04 (49)	1.70 ± 0.22 (18)	0.34 ± 0.04 (112)	0.10 ± 0.01 (159)
layer 0.35-1.40 nm	2.18 ± 0.11 (809)	2.22 ± 0.08 (1184)	2.26 ± 0.11 (956)	2.07 ± 0.12 (779)	1.10 ± 0.08 (1020)	0.38 ± 0.04 (1468)
all	2.48 ± 0.03	2.38 ± 0.01	2.42 ± 0.00	2.52 ± 0.00	1.31 ± 0.02	0.44 ± 0.00
SPC						
layer 0-0.35 nm	2.32 ± 0.04 (17)	1.61 ± 0.05 (43)	2.11 ± 0.28 (38)	3.96 ± 0.11 (10)	0.64 ± 0.05 (66)	0.28 ± 0.04 (82)
layer 0.35-1.40 nm	4.28 ± 0.05 (747)	4.09 ± 0.15 (989)	4.01 ± 0.08 (811)	4.32 ± 0.01 (688)	2.51 ± 0.02 (919)	1.21 ± 0.05 (996)
all	4.21 ± 0.02	4.04 ± 0.04	4.11 ± 0.02	4.20 ± 0.02	2.68 ± 0.00	1.36 ± 0.01

Comparison of reduced point charge models of proteins: Molecular Dynamics simulations of Ubiquitin

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SI 1 Computational details

Search for critical points in a smoothed molecular property

As already reported before [31], we adapted Leung *et al.*'s idea [41] to 3D molecular property functions. The various steps of the resulting merging/clustering algorithm are as follows. First, at scale $s = 0$, each atom of a molecular structure is considered either as a local maximum (peak) or minimum (pit) of the scalar field f . All atoms are consequently taken as the starting points of the merging procedure. Second, as s increases from 0 to a given maximal value s_{max} , each point moves continuously along a gradient path to reach a location in the 3D space where $\nabla f(s) = 0$. On a practical point of view, this consists in following the trajectory of the peaks and pits within the molecular property calculated at s . The trajectory search is stopped when $|\nabla f(s)|$ is lower or equal to a limit value, $grad_{lim}$. Once all peak/pit locations are found, close points are merged if their inter-distance is lower than the initial value of $\Delta^{1/2}$. The procedure is repeated for each selected value of s . If the initial Δ value is too small to allow convergence towards a local maximum or minimum within the given number of iterations, its value is doubled (a scaling factor that is arbitrarily selected) and the procedure is repeated until final convergence.

In the present work, to design amino acid point charge models, extrema were located using the merging/clustering algorithm applied to the CD distribution functions smoothed at $s = 1.7$ bohr², with $\Delta_{init} = 10^{-4}$ bohr² and $grad_{lim} = 10^{-6}$ e⁻ bohr⁻², and to the PASA ED distribution functions smoothed at $s = 1.4$ bohr², with $\Delta_{init} = 10^{-4}$ bohr² and $grad_{lim} = 10^{-5}$ e⁻ bohr⁻².

Charge calculation

To stay consistent with the analytical expression of the Amber99 FF, only point charge values are assigned to each of the CPs of a 3D molecular property field. The charge fitting program QFIT [Borodin O, Smith GD. Force Field Fitting Toolkit, The University of Utah] was used as detailed in ref. [31]. All MEP grids were built using the Amber99 [37] atomic charges which were assigned using the software PDB2PQR [42]. Side chains and main chains of the AAs were treated separately, as discussed in ref. [31]. In all fittings, the total electric charge and the magnitude of the molecular dipole moment were constrained to be equal to the corresponding all-atom Amber99 values.

Free energy calculations using the Bennett acceptance ratio (BAR)

To allow the calculation of free energy differences, the Bennett's acceptance ratio (BAR) [45,46] is based on the statistical mechanics expression:

$$\Delta F_{0 \rightarrow 1} = \beta^{-1} \ln \frac{Z_0}{Z_1} \quad (1)$$

where $\beta^{-1} = kT$. Z_i is the configurational integral, *i.e.*, the contribution to the partition function that involves the interaction energy U_i between the particles of a system. If one introduces a weight function W , the ratio of configurational integrals can be written as:

$$\frac{Z_0}{Z_1} = \frac{\langle W e^{-\beta U_0} \rangle_1}{\langle W e^{-\beta U_1} \rangle_0} \quad (2)$$

Bennett showed that a suitable choice for W , which minimizes the square error on $\Delta F_{0 \rightarrow 1}$, is:

$$W = C \left(\frac{Z_0}{n_0} e^{-\beta U_1} + \frac{Z_1}{n_1} e^{-\beta U_0} \right)^{-1} \quad (3)$$

where, in practice, n_i is the number of configurations sampled for state i , and:

$$C = \frac{1}{\beta} \ln \frac{Z_0 n_1}{Z_1 n_0} \tag{4}$$

When multiple intermediate states are used, an estimation of the total free energy difference $\Delta F_{0 \rightarrow 1}$ is obtained as a summation over free energy differences between the successive intermediate states.

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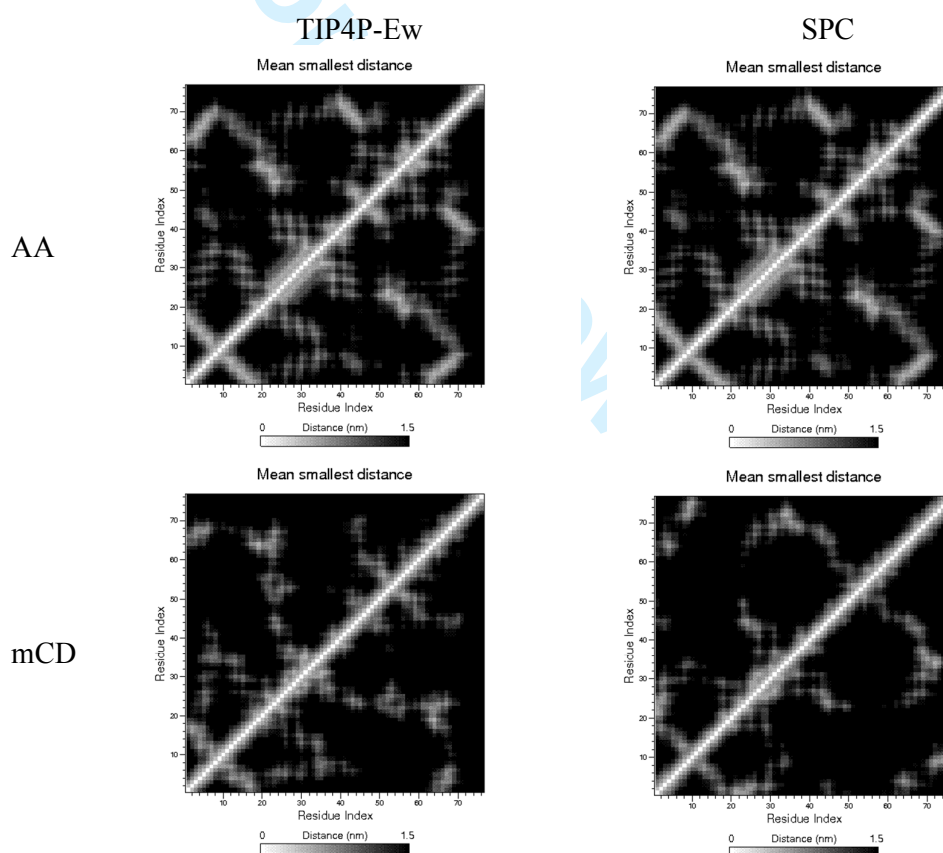
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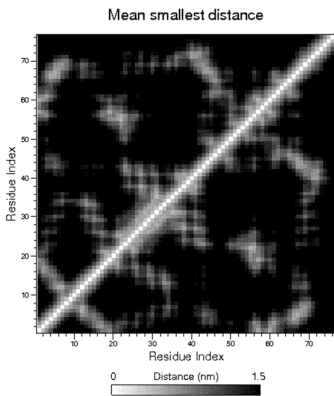
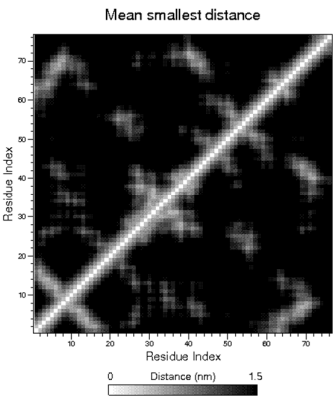
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SI 2 C_{α} - C_{α} distance maps of Ubiquitin calculated for the last frame of the Amber99SB-based 20 ns MD trajectories in water at 300 K, obtained using the all-atom, mCD, mCDa, and mPASA models.

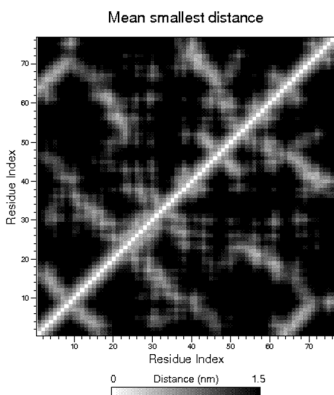
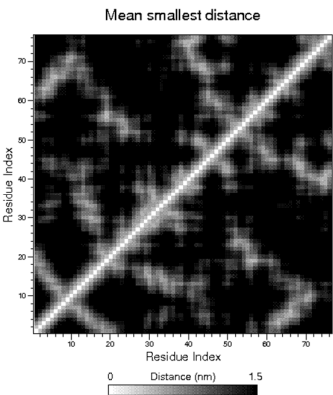


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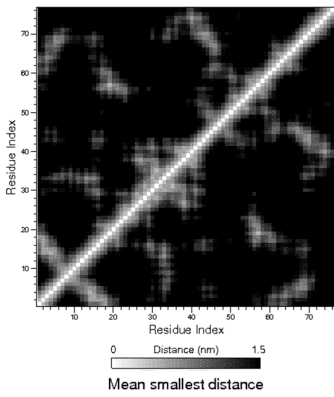
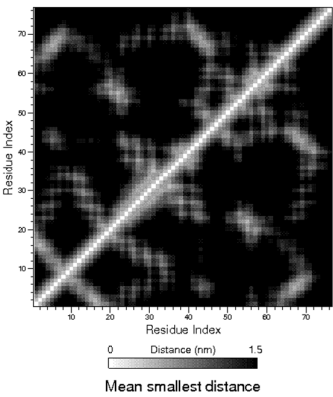
mCDa



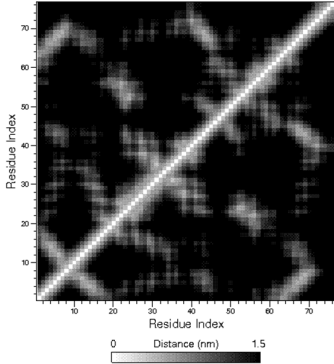
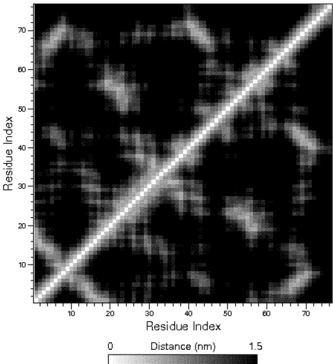
mPASA



mCD
(277 K)



mCD
(250 K)



Comparison of reduced point charge models of proteins: Molecular Dynamics simulations of Ubiquitin

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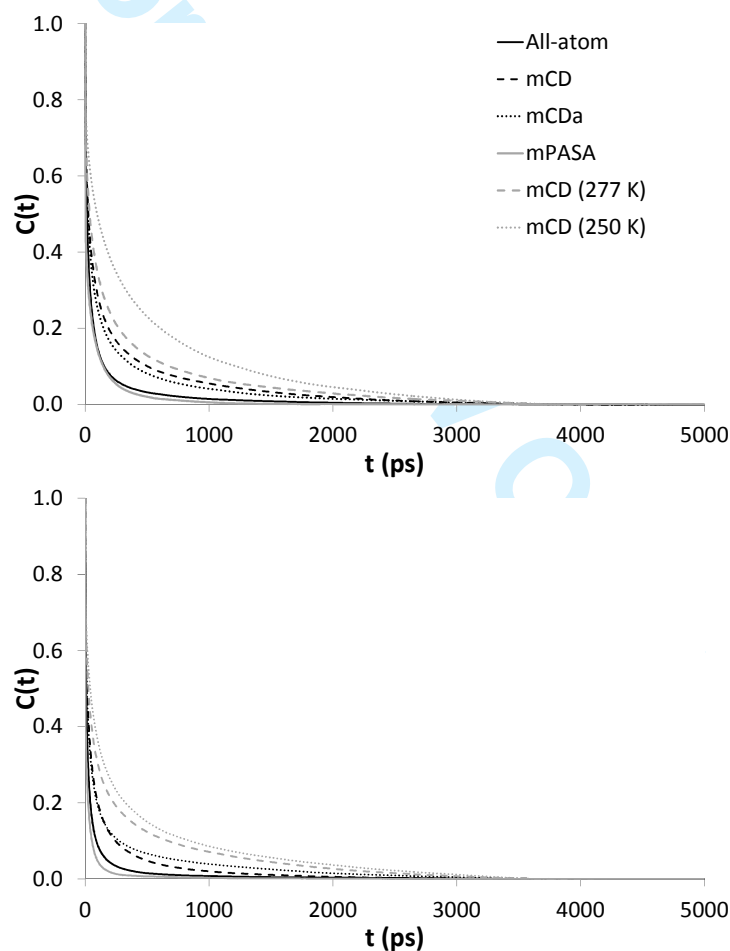
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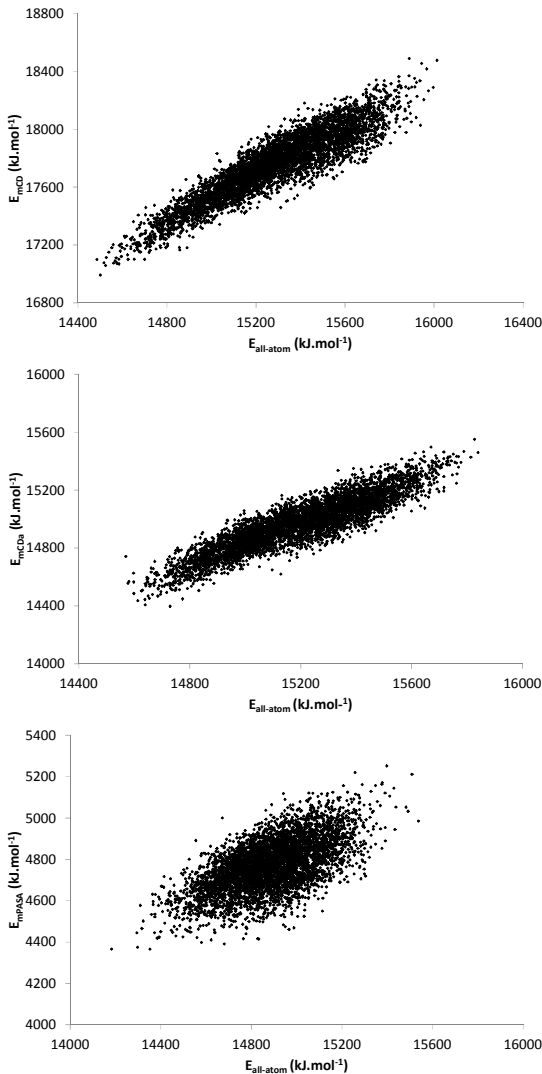
SI 3 Autocorrelation functions of Ubiquitin-water H-bonds calculated from the last 10 ns of the Amber99SB-based MD trajectories at 300 K, obtained using the all-atom, mCD, mCDa, and mPASA models with the (Top) TIP4P-Ew and (Bottom) SPC FFs.



Comparison of reduced point charge models of proteins:
Molecular Dynamics simulations of Ubiquitin

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SI 4 Correlation between the all-atom and RPCM potential energy values of Ubiquitin calculated from the 20 ns MD trajectories using the TIP4P-Ew and AMBER99SB force fields at 300 K. (Top) E_{mCD} versus $E_{\text{all-atom}}$, (Middle) E_{mCDa} versus $E_{\text{all-atom}}$, and (Bottom) E_{mPASA} versus $E_{\text{all-atom}}$.



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Comparison of reduced point charge models of proteins: Molecular Dynamics simulations of Ubiquitin

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SI 5 Energy values of the optimized structures of Ubiquitin (kJ.mol⁻¹) and averages with standard deviation values calculated over the last 10 ns of the Amber99SB-based MD simulations of Ubiquitin in TIP4P-Ew and SPC water, at T = 300 K, obtained using the all-atom, mCD, mCDa, and mPASA models. Energy terms such as short-range Coulomb (Cb-SR), short- and long-range Lennard-Jones (LJ-SR and LJ-LR), 1-4 Coulomb (Cb-14), and 1-4 Lennard-Jones (LJ-14) were calculated during a post-processing stage. Values in italics are for the SPC water model.

	Optimized structure				Solvated structure TIP4P-Ew SPC				Vacuum			
	All-atom	mCD	mCDa	mPASA	All-atom	mCD	mCDa	mPASA	All-atom	mCD	mCDa	mPASA
Intra-molecular												
Stretching	137.42	96.05	90.63	81.41	929.34 ± 51.48 <i>928.93 ± 51.17</i>	904.70 ± 51.10 <i>894.95 ± 49.53</i>	886.02 ± 49.14 <i>881.42 ± 49.70</i>	869.87 ± 49.83 <i>870.13 ± 49.09</i>	730.29 ± 40.02	891.19 ± 48.85	883.75 ± 48.92	883.14 ± 49.62
Bending	682.21	644.85	627.05	567.21	2536.24 ± 80.95 <i>2546.24 ± 78.59</i>	2466.70 ± 79.47 <i>2428.38 ± 74.50</i>	2427.25 ± 75.91 <i>2420.10 ± 75.58</i>	2449.21 ± 77.60 <i>2432.35 ± 75.98</i>	2129.09 ± 65.64	2598.66 ± 75.62	2563.25 ± 77.15	2503.71 ± 76.77
Dihedral	3058.19	2983.23	2994.27	2953.83	3123.74 ± 39.02 <i>3126.55 ± 40.05</i>	3039.97 ± 40.14 <i>3074.94 ± 40.51</i>	3083.32 ± 40.43 <i>3069.35 ± 37.97</i>	3040.52 ± 39.77 <i>3064.81 ± 41.97</i>	3145.13 ± 32.21	3360.80 ± 39.31	3313.17 ± 42.09	3198.14 ± 44.99
Improper-Dihedral	50.40	48.33	39.76	20.64	149.45 ± 16.28 <i>149.87 ± 16.43</i>	134.46 ± 14.78 <i>134.73 ± 15.20</i>	135.19 ± 14.67 <i>135.29 ± 15.24</i>	130.33 ± 14.25 <i>131.97 ± 14.18</i>	129.63 ± 12.82	165.87 ± 17.38	156.86 ± 16.46	142.25 ± 15.21
Cb-SR	-4461.87	-3944.84	-3464.36	-99.70	-4517.21 ± 124.45 <i>-4613.55 ± 122.58</i>	-3841.82 ± 179.00 <i>-4355.43 ± 135.40</i>	-3334.40 ± 114.63 <i>-3592.77 ± 113.68</i>	-504.91 ± 77.36 <i>-418.16 ± 78.58</i>	-6831.93 ± 62.47	-6572.02 ± 67.42	-5915.33 ± 83.30	-1478.09 ± 34.70
LJ-SR	-2305.34	-2257.15	-2262.03	-2387.91	-2203.11 ± 41.70 <i>-2227.90 ± 45.82</i>	-1531.84 ± 52.64 <i>-1446.46 ± 92.23</i>	-1685.80 ± 79.60 <i>-1879.23 ± 46.29</i>	-2342.62 ± 49.54 <i>-2386.84 ± 49.62</i>	-2099.38 ± 44.18	-1933.72 ± 54.69	-2041.19 ± 53.84	-2215.05 ± 52.32
LJ-LR	-57.71	-57.61	-57.69	-57.71	-56.56 ± 0.51 <i>-56.87 ± 0.53</i>	-41.28 ± 0.87 <i>-39.70 ± 1.91</i>	-43.59 ± 2.51 <i>-50.51 ± 0.80</i>	-57.80 ± 0.79 <i>-57.31 ± 0.79</i>	-60.04 ± 0.33	-59.09 ± 0.41	-59.96 ± 0.50	-58.30 ± 0.55
Cb-14	12463.77	15648.03	12465.32	0	12272.70 ± 86.70 <i>12317.49 ± 90.13</i>	15564.52 ± 78.75 <i>15808.82 ± 75.25</i>	12434.40 ± 75.91 <i>12501.92 ± 70.03</i>	0	12290.70 ± 64.67	15672.66 ± 54.15	12667.13 ± 60.77	0
LJ-14	1193.46	1272.22	1274.89	1196.27	1139.32 ± 29.10 <i>1142.58 ± 28.58</i>	1174.64 ± 30.44 <i>1168.45 ± 30.60</i>	1171.29 ± 30.74 <i>1193.81 ± 31.55</i>	1174.53 ± 32.00 <i>1164.35 ± 30.96</i>	1082.14 ± 23.01	1204.27 ± 31.57	1196.19 ± 31.13	1164.11 ± 31.45
Inter-molecular												
Cb-SR					-8024.49 ± 259.24 <i>-7443.38 ± 252.90</i>	-12127.84 ± 442.91 <i>-10811.59 ± 322.04</i>	-11436.43 ± 292.18 <i>-9923.32 ± 285.76</i>	-3423.91 ± 201.87 <i>-3579.74 ± 200.73</i>				
LJ-SR					-506.14 ± 79.05 <i>-512.46 ± 78.07</i>	-708.75 ± 105.24 <i>-895.29 ± 120.34</i>	-645.90 ± 132.51 <i>-585.34 ± 94.15</i>	-1026.21 ± 65.66 <i>-949.90 ± 58.61</i>				
LJ-LR					-109.38 ± 0.84 <i>-104.94 ± 0.87</i>	-127.61 ± 1.08 <i>-124.18 ± 2.12</i>	-124.13 ± 3.01 <i>-112.51 ± 1.08</i>	-104.06 ± 1.05 <i>-100.86 ± 1.06</i>				

	Solvated structure TIP4P-Ew SPC		Vacuum	
	mCD (277 K)	mCD (250 K)	mCD (277 K)	mCD (250 K)
Intra-molecular				
Stretching	840.39 ± 47.56 842.23 ± 46.99	771.22 ± 42.99 764.70 ± 42.63	896.08 ± 50.39	825.65 ± 44.92
Bending	2302.58 ± 70.44 2304.26 ± 70.64	2124.88 ± 64.14 2102.81 ± 67.14	2568.14 ± 76.23	2376.84 ± 68.79
Dihedral	30413.57 ± 35.64 3025.77 ± 39.21	3012.22 ± 33.72 3032.66 ± 32.37	3243.14 ± 36.99	3274.06 ± 35.66
i-Dihedral	128.97 ± 14.22 125.44 ± 13.83	118.79 ± 13.18 117.27 ± 13.11	159.71 ± 16.98	155.89 ± 15.16
Cb-SR	-4079.09 ± 139.23 -3901.70 ± 121.85	-4271.52 ± 123.21 -4379.75 ± 111.77	-6639.78 ± 69.98	-6673.25 ± 69.10
LJ-SR	-1820.78 ± 41.30 -1622.03 ± 49.38	-1998.39 ± 43.54 -1967.53 ± 39.42	-2032.61 ± 53.70	-2050.94 ± 54.35
LJ-LR	-49.71 ± 0.70 -44.51 ± 1.30	-51.67 ± 0.62 -51.15 ± 0.49	-58.91 ± 0.38	-59.59 ± 0.39
Cb-14	15751.50 ± 66.35 15560.32 ± 82.55	15772.30 ± 59.09 15743.36 ± 63.66	12115.84 ± 79.16	12200.61 ± 75.46
LJ-14	1162.37 ± 258.56 1170.26 ± 30.50	1195.60 ± 28.22 1156.78 ± 27.17	1126.68 ± 27.79	1102.38 ± 25.31
Inter-molecular				
Cb-SR	-11559.06 ± 313.70 -11444.40 ± 295.59	-10763.32 ± 276.88 -9995.22 ± 254.60		
LJ-SR	-551.29 ± 96.11 -711.68 ± 96.37	-454.20 ± 83.77 -557.58 ± 83.48		
LJ-LR	-120.05 ± 0.82 -121.46 ± 1.58	-117.11 ± 0.76 -115.71 ± 0.79		